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(71) Applicant (for all designated States except US): NATIONAL RESEARCH DEVELOPMENT CORPORATION [GB/GB]; 101 Newington Causeway, London SE1 6BU (GB).

(72) Inventors; and
(75) Inventors/Applicants (for US only): PICKUP, John, Christopher [GB/GB]; Weavers End, Course Horn Lane, Cranbrook, Kent TN17 3NR (GB). CLAREMONT, Denzil, Joseph [GB/GB]; 6 Prentice Court, Leopold Avenue, Wimbledon, London SW19 7HA (GB).

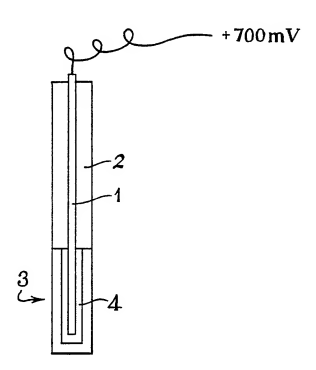
(74) Agent: NEVILLE, Peter, Warwick; Patent Department, National Research Development Corporation, 101 Newington Causeway, London SE1 6BU (GB).

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(54) Title: GLUCOSE-SENSING ELECTRODE



(57) Abstract

A glucose sensing electrode consists of a platinum wire (1) coated in synthetic hydrophilic polyurethane or polyhydroxyethylmethacrylate (4) containing glucose oxidase, this being itself coated in a mixture of hydrophobic polyurethane and hydrophilic polyhydroxyethylmethacrylate.

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GLUCOSE-SENSING ELECTRODE

FIELD OF THE INVENTION

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This invention relates to a glucose-sensing electrode and to a glucose sensor using it. Such a sensor may find use in single measurements in, for example, samples of body fluids outside the body (e.g. blood, serum, plasma, urine), or, as an implantable sensor, in continuous measurement of glucose concentrations within the body.

DESCRIPTION OF PRIOR ART

Glucose sensors are probes which ideally would give a rapid and specific signal proportional to glucose concentration, without the need for added reagent. They have major potential applications in the management of diabetes mellitus, for home blood glucose monitoring (where a capillary blood sample obtained by finger-prick is spotted onto the sensor), for improved laboratory and ward glucose analysers which are expensive and bulky, and as implantable devices for continuous measurement of body glucose levels. The latter could be used as a read-out of diabetic "control" or to trigger an alarm when glucose concentrations fall to dangerously low or high levels (hypoand hyperglycaemia). Alternatively, an implantable glucose sensor could be linked to an insulin infusion pump to provide automatic feedback control of insulin delivery (an artificial endocrine pancreas).

Other uses for glucose sensors in non-medical areas include monitoring or fermentation processes and food analysis.

Several technologies for glucose sensors have been described, including amperometric and potentiometric enzyme electrodes and optical approaches. For example, Diabetes Research Clinical Practice Supplement 1985 Vol 1 page S447 item 1162 states "Amperometric glucose sensors which are relatively oxygen insensitive have been constructed using 1,1'-dimethyl ferrocene (dicyclopentadienyl iron) to mediate glucose oxidase catalysed electron transfer between glucose and a carbon (graphite) base electrode. Several 1mm wide probe sensors, mounted on Plexiglass and suitable for in vivo implantation, were tested simultaneously

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in vitro using a computer-assisted apparatus which poises the working electrode potential at +160 mV and records the current output. Electrodes dip-coated in 4% polyurethane were generally linear to at least 20 mmol/1 glucose with a 95% response time of 05 30 sec. to 3 min." It was known, e.g. from Updike et al, Diabetes Care 1982 Vol 5 page 209 that "Oxygen is a hydrophobic gas and thus diffuses best through a hydrophobic membrane. the other hand, glucose is a hydrophilic substance and diffuses best through a hydrophilic membrane", these being applied between the electrode and a layer of glucose oxidase. It is consequently 10 taught by Updike to extend the linear range of glucose response by adding an outer hydrophobic layer to the glucose oxidase. Shichiri et al, The Lancet 20 November 1982 p 1129, describe a glucose sensor in which a platinum anode is coated with glucose oxidase immobilised in heparin and cellulose diacetate, an outer layer of polyurethane then being applied. The major problems with most sensors to date are unpredictable drift in output, sensitivity to changes in oxygen concentration at the sensing site and complex or intricate manufacturing procedures unsuitable for mass production.

DESCRIPTION OF THE INVENTION

According to the present invention, a glucose-sensing comprises platinum (e.g. wire), or electrode another base electrode capable electrochemically of oxidising hydrogen peroxide. coated in hydrophilic matter containing oxidase, characterised in that said matter is a synthetic hydrophilic polymer. Preferably the polymer is applied in predominantly ethanolic aqueous solution, this having been found to be minimally damaging to the activity of the glucose oxidase enzyme compared with acetone, which has previously been used for coating enzyme-containing polymer. The polymer is preferably hydrophilic polyurethane and/or polyhydroxyethylmethacrylate.

Preferably, the coating of hydrophilic matter is itself coated in a mixture of hydrophobic matter and hydrophilic matter,

as we have found that this may extend the linearity of response of the electrode to glucose. This mixture may be polymeric; the hydrophobic component may comprise hydrophobic polyurethane, and the hydrophilic component is hydrophilic polyurethane and/or polyhydroxyethylmethacrylate.

The invention extends to a glucose sensor comprising the electrode set forth above.

The invention will now be described by way of example with reference to the accompanying drawings.

10 BRIEF DESCRIPTION OF THE DRAWINGS

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Figure 1 shows a glucose-sensing electrode according to the invention,

Figure 2 shows the current passed by two electrodes according to the invention at varying glucose concentrations in vitro,

Figure 3 shows the mean current at various times, and standard error of the mean, for a group of five electrodes according to the invention operated in vitro for 24 hours, this to illustrate the stability of the electrodes,

Figure 4 shows the in vivo response of an electrode according to the invention compared with the blood glucose level analysed conventionally, and

Figure 5 shows the effect of varying oxygen concentration at fixed glucose concentration on the output of electrodes according to the invention.

25 EXAMPLE OF THE INVENTION

Turning to Figure 1, the construction of a glucose-sensing electrode according to the invention is shown.

Platinum wire 1 is dipped in insulating varnish 2 and baked at 80°C for 2hr in an oven. After cooling, an approximately 5mm length 3 is scraped from one end of the wire for application of sensing material 4. This material consists of a mixture of (a) glucose oxidase enzyme dissolved in water and (b) either polyhydroxyethylmethacrylate polymer or hydrophilic polyurethane in 75% ethanol/25% water. The concentration of (a) is 1000 IU/ml, and of (b) is 12.5% w/v, and the volume ratio of (a):(b)

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is 1:3. Alternatively, (a) as a step in making the mixture can be omitted, and dry glucose oxidase mixed with the hydrophilic polymer in the ethanol/water to a total concentration of say 5000 IU/ml glucose oxidase. After dip-coating in the enzyme/polymer mixture, the wire is air-dried at room temperature for 20 min. This is simpler than covalent enzyme-attachment procedures using substances such as glutaraldehyde.

An outer membrane application material consists of a mixture of (a) hydrophobic polyurethane in tetrahydrofuran optionally with ethanol (concentration 10%) and (b) either the hydrophilic polymer polyhydroxyethylmethacrylate in 75% ethanol or hydrophilic polyurethane in tetrahydrofuran, concentration 10%, ratio a:b 3:1. After dip-coating in this material, the electrode is air dried for 20 min at room temperature.

In vitro, a pseudoreference (combined reference and auxiliary electrode) consisting of silver/silver chloride adjacent to the working electrode described, or in vivo a silver/silver chloride electrocardiogram electrode on the skin above the implanted working electrode, makes a glucose sensor according to the invention. The working electrode is operated in amperometric mode at a fixed 700mV.

Before routine use, the sensor may be preconditioned by operating at +700mV in 5mmol/l buffered glucose solution at pH 7.4 for 18 hr at $37^{\circ}C$.

Fig. 2 shows in vitro calibration curves for two glucose sensors, which demonstrate high current outputs which increase linearly to at least 20 mmol/l glucose. These sensors were as described for Figure 1, with 10% polyhydroxyethyl-methacrylate/glucose oxidase inner membrane and 30 75% hydrophobic polyurethane/25% hydrophilic polyhydroxyethyl-methacrylate outer membrane.

Fig. 3 shows the mean current \pm standard error of the mean for a group of 5 glucose sensors operated <u>in vitro</u> at 37°C for 24hr in 5mmol/l glucose solution, which demonstrates the excellent stability of the devices.

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Fig. 4 shows the in vivo response of an electrode according to the invention compared with the blood glucose level analysed conventionally. This demonstrates the possibility of the use of the glucose sensor as an implantable monitor of glycaemic control in man. A sensor was implanted in the subcutaneous tissue of the forearm of a normal volunteer subject. The potential was set at +700mV using a potentiostat and a surface electrocardiogram electrode used as a reference in a two-electrode configuration. At time 0, 75g of glucose was administered orally and blood glucose concentrations measured conventionally using a Yellow Springs Instruments analyser, a bench-top instrument for in vitro blood sample analysis, having an immobilised glucose oxidase membrane and operating by detection of the produced hydrogen peroxide. Electrode current readings were calibrated by assuming that the initial current value is equivalent to the initial blood glucose level. The graph demonstrates that tissue glucose levels measured by the sensor according to the invention follow blood glucose levels (measured conventionally) with little delay, though of lesser magnitude. The implanted sensor can thus be used to monitor changes in blood glucose levels.

Fig. 5 shows an example of the mean current output of 5 glucose sensors operated in a fixed concentration of buffered glucose solution in vitro at 20 kPa $p[O_2]$ and at 1 kPa $p[O_2]$ (viz, one-fifth of an atmosphere of oxygen (as normal) and 0.01 atmosphere respectively) which demonstrates that the sensors are not significantly affected by changing concentrations of oxygen. These sensors did not have the outer mixed-hydrophobic-hydrophilic membrane.

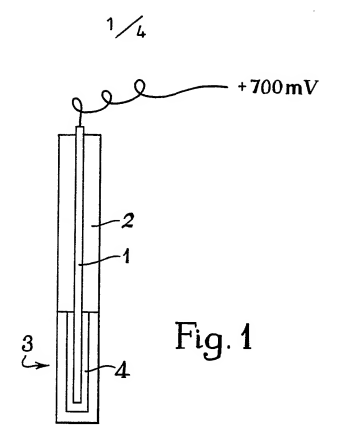
Thus, such a glucose sensor could be used as an implantable 30 device:

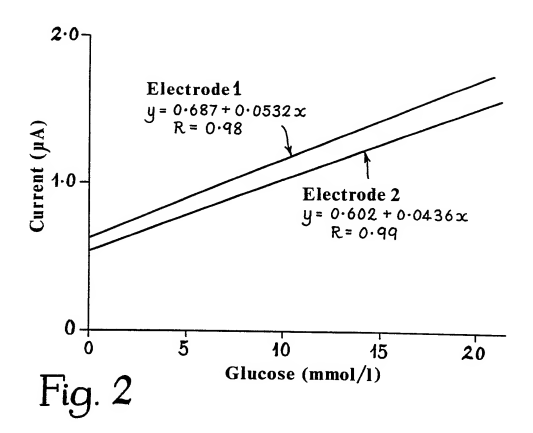
- (a) to give a continuous measure of glycaemic control.
- (b) to detect hypoglycaemia and sound an alarm at a pre-set glucose level
- (c) to detect hyperglycaemia and sound an alarm at a pre-set glucose level

- (d) to be linked to a portable or implanted insulin infusion pump with feedback control of insulin delivery according to the prevailing glucose levels and also as a non-implanted device:
- (e) to provide a small, hand-held device for monitoring glucose levels in blood, serum or plasma samples or in other body fluids. This might be configured as a pen-type device with digital readout and incorporating the sensor at one end.
 - (f) as a small bench-top glucose analyser for laboratory, ward, bed-side, office or other use, and
- 10 (g) to form part of a multi-analyser system (glucose plus other analytes).

CLAIMS

- 1. A glucose-sensing electrode, comprising an inert conductor coated in hydrophilic matter containing glucose oxidase, characterised in that said matter is a synthetic hydrophilic polymer.
- 05 2. An electrode according to Claim 1, wherein said polymer is applied in predominantly ethanolic and/or aqueous solution.
 - 3. An electrode according to Claim 1, wherein the polymer is hydrophilic polyurethane and/or polyhydroxyethylmethacrylate.
- 4. An electrode according to Claim 2 wherein the polymer is hydrophilic polyurethane and/or polyhydroxyethylmethacrylate.
 - 5. An electrode according to Claim 1, 2, 3 or 4 wherein the coating of hydrophilic matter is iself coated in a mixture of hydrophobic matter and hydrophilic matter.
- 6. An electrode according to Claim 5, wherein the mixture is polymeric.
 - 7. An electrode according to Claim 5, wherein the hydrophobic matter comprises hydrophobic polyurethane.
 - 8. An electrode according to Claim 6 wherein the hydrophobic matter comprises hydrophobic polyurethane.
- 9. An electrode according to Claim 6, 7 or 8 wherein the hydrophilic matter of said mixture comprises hydrophilic polyurethane and/or polyhydroxyethylmethacrylate.
 - 10. An electrode according to Claim 1, 2, 3, 4, 6, 7 or 8 wherein the inert conductor is platinum.
- 25 11. An electrode according to Claim 5 wherein the inert conductor is platinum.
 - 12. An electrode according to Claim 9 wherein the inert conductor is platinum.





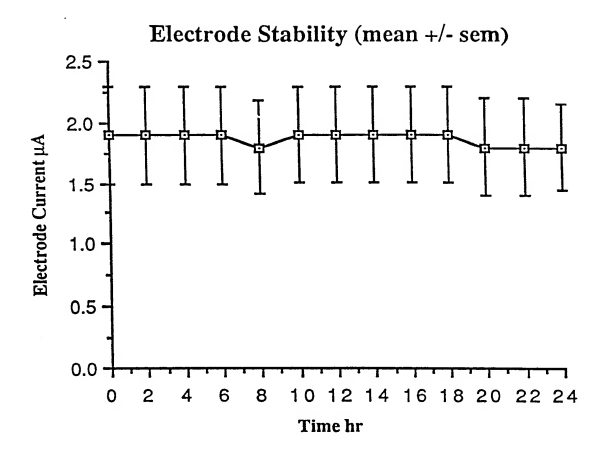


Fig. 3

Blood and Tissue Glucose Concentrations

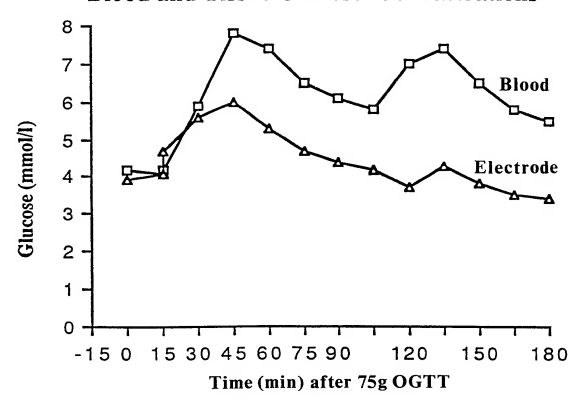


Fig.4

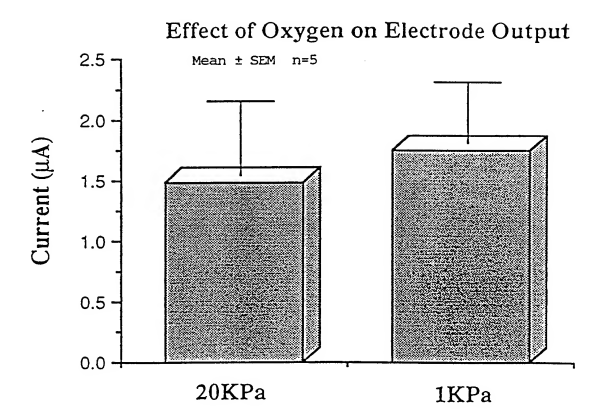


Fig. 5

International Application N

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III. DOCUME	NTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET)	PCT/GB 90/00660
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9000660

SA 36677 This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information. 14/0 14/08/90

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